**Introductory nanoscience:**

*Physical and chemical concepts*

Masaru Kuno

Garland Science: Taylor & Francis Group, 2011

420 pages, $115


**Hume-Rothery rules for structurally complex alloy phases**

Uichiro Mizutani

CRC Press, 2011

342 pages, $89.95 (pay per view)

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**Hume-Rothery’s work in the 1920s and 1930s turned the art of metallurgy into science and contributed to the emergence of solid-state physics.**

Hume-Rothery or H-R rules express the conditions for alloy formation in terms of the difference in atomic diameters of the constituent elements, electronegativity difference, and electron concentration (electron/atom) ratio of the alloy. Hume-Rothery (H-R) rules for formation of alloys continue to be important. For example, quasicrystals, whose crystallographic structures did not follow the classical definition of crystals, respect these rules and some of them were discovered using H-R rules as guidelines. In this book by Uichiro Mizutani, the main concern is the electron concentration rule and the book provides deep insights into this topic using first-principles band calculations.

The book is well organized. The first three chapters review the early theories. Several key concepts are discussed in chapters 4 to 6, such as mechanisms for the formation of pseudogap at the Fermi level, using linear muffin-tin orbital-atomic-sphere approximation (LMTO-ASA), and the full-potential linearized augmented plane wave (FLAPW). This part of the book assumes a background in quantum mechanics. The calculations are made on alloys containing 52 atoms per unit cell, whereas the term “structurally complex alloys” used in the title of the book would include giant unit cells with thousands of atoms with well-defined clusters. Chapters 7 to 10 apply the concepts developed to metallic alloys. The author introduces “the effective Fermi sphere” and H-R plots based on first-principles calculations. Mizutani derives an effective valency for transition metals (TMs) and points out they take positive values which depend on the atomic environment. Frequently, negative values suggested by Raynor for TM are used in calculations for Al-TM alloys and positive values suggested by H-R for the same TM in the case of Cu–TM alloys. Mizutani shows that the use of valence electron concentration instead of valency assigned in the periodic table results in a better correlation with many properties. For example, Cu has a value of 11 and 1, respectively, for these parameters.

In summary, this book will serve as a good reference work for graduate students in metallurgy and physics and researchers in metallurgical laboratories and industries. It is Hume-Rothery revisited in the light of first-principles calculations. It is therefore appropriate to point out one feature of the concluding chapter 11. Hume-Rothery’s famous book in 1948, *Electrons, Atoms, Metals and Alloys*, is entirely in the form of a conversation between a young scientist and an old metallurgist. Mizutani adopts the same format for the last chapter of this book. The dialogue continues.

**Reviewer:** N. Balasubramanian

works at the Center for Science, Technology and Policy, Bangalore, India.
Instead of being encyclopedic in coverage, the book does an excellent job in fleshing out details often omitted in more standard textbooks. It carefully points out pitfalls in notation, for instance, noting that the [001] direction need not be perpendicular to the (001) plane for all crystal classes and noting the ambiguities in ways to report Einstein A and B coefficients. There is a relatively complete derivation of the Scherrer equation for estimating crystallite size from x-ray diffraction measurements in contrast to most texts which simply state the end result. The Kronig-Penney model for a one-dimensional periodic crystal is also described thoroughly; students less adept at linear algebra should be able to more clearly follow through the steps to the final expression for energies. At various points, the text also provides back-of-the-envelope calculations; the book explains in detail the spring model commonly used to understand frequency shifts in Fourier Transform infrared spectroscopy spectra. 

Introductory Nanoscience is structured like a standard text with chapter summaries, problems, and references for each chapter. The problems differ significantly in nature and difficulty from chapter to chapter. Some point to interesting results from the current literature worthy of further study, a number involve straightforward numerical substitutions while still others are relatively standard analytical problems found in traditional quantum mechanics texts. Overall, the reference sections are generous and should guide students to other resources as needed.

In summary, Introductory Nanoscience provides students entering the field with more details and explanations in comparison to other sources. As such, the book should be of value to students, although due to its lack of coverage in certain areas and unevenness in the problems, this text may not be broad or balanced enough to serve as the single required book in a course on nanoscience.

Reviewer: Yumi Ijiri is a professor of physics in the Department of Physics and Astronomy at Oberlin College.